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NMR Identification of the Binding Surfaces Involved in the Salmonella and Shigella Type III Secretion Tip-Translocon Protein-Protein Interactions

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Abstract

The type III secretion system (T3SS) is essential for the pathogenesis of many bacteria including *Salmonella* and *Shigella*, which together are responsible for millions of deaths worldwide each year. The structural component of the T3SS consists of the needle apparatus, which is assembled in part by the protein-protein interaction between the tip and the translocon. The atomic detail of the interaction between the tip and the translocon proteins is currently unknown. Here, we used NMR methods to identify that the N-terminal domain of the *Salmonella* SipB translocon protein interacts with the SipD tip protein at a surface at the distal region of the tip formed by the mixed a/β domain and a portion of its coiled-coil domain. Likewise, the *Shigella* IpaB translocon protein and the IpaD tip protein interact with each other using similar surfaces identified for the *Salmonella* homologs. Furthermore, removal of the extreme N-terminal residues of the translocon protein, previously thought to be important for the interaction, had little change on the binding surface. Finally, mutations at the binding surface of SipD reduced invasion of *Salmonella* into human intestinal epithelial cells. Together, these results reveal the binding surfaces involved in the tip-translocon protein protein interaction and advance our understanding of the assembly of the T3SS needle apparatus.

Keywords

type III secretion system; tip; translocon; SipB; IpaB; SipD; IpaD; NMR spectroscopy

INTRODUCTION

Many Gram-negative bacteria assemble a protein nanoinjector called the type III secretion system (T3SS) to inject virulence factors into the cytoplasm of host cells and initiate infectious diseases. This includes the causative agents of food poisoning/typhoid (*Salmonella Typhimurium/Typhi*) and dysentery (*Shigella flexneri*). Together, these pathogens are responsible for millions of deaths worldwide each year, primarily in children

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under the age of six.^{1,2} The T3SS is essential for pathogenicity as deletions of its protein components render bacteria non-invasive.³ The structural component of the T3SS, the needle apparatus, is composed of a base anchored at the bacterial cell membranes, an extracellular needle, a tip complex and a translocon.⁴ The needle is formed by the oligomerization of PrgI in *Salmonella* and MxiH in *Shigella* and the atomic structure of the polymerized needle has been determined for both *Salmonella* and *Shigella*.^{5,6} At the needle tip sits a complex of an estimated pentameric copy of SipD in *Salmonella*⁷ and IpaD in *Shigella*.^{8,9} The tip complex functions as a signal transducer for host cell contact, a platform for the assembly of the translocon, and a regulator of secretion of effector proteins.^{7,10,11} The crystal structures of SipD^{12,13} and IpaD^{14,15} reveal three domains: an N-terminal α -helical hairpin, a central coiled-coil and a mixed α/β region (Fig. 1). The N-terminal α -helical hairpin functions as a self-chaperone that prevents self-oligomerization and premature interactions with other T3SS proteins, such as the needle.^{15,16} A current hypothesis is that during assembly at the tip of the needle, the N-terminal α -helical hairpin of SipD/IpaD is displaced to allow interaction with the needle.^{15,16}

Upon host cell contact, bacteria assemble a ~3 nm wide translocon pore in the host membrane to allow the passage of effectors into the host cytoplasm.^{17–19} The translocon is assembled from two transmembrane proteins termed the major and the minor translocon protein based on their molecular weights. The major and minor translocon proteins in *Salmonella* are SipB and SipC, respectively; and their counterparts in *Shigella* are IpaB and IpaC, respectively.^{3,4,17} The translocon proteins also function as effectors.^{20–22} The topology of SipB and IpaB are predicted to contain an N-terminal ectodomain, a transmembrane helical region, and a C-terminal amphipathic helix.^{23–27} The atomic structure of the assembled translocon or the full length translocon proteins is currently unknown, however, crystal structures of the N-terminal ectodomains of SipB and IpaB revealed structurally similar coiled-coil motifs.²⁸ Further, Nguyen *et al.*²⁹ recently proposed a crystallography-derived model of the membrane-insertion and topology of the SipB/IpaB counterpart in *Aeromonas hydrophila*, AopB.

How the translocon proteins interact with the tip protein is poorly understood. Dickenson et al.23 reported the interaction between the Shigella IpaD tip protein with the N-terminal ectodomain of IpaB (residues 11-226) by FRET and showed that the interaction required the presence of the bile salt deoxycholate and that IpaB residues 11-27 were essential for the interaction. Results showing that the proximal region of SipD (defined here as the bottom of the coiled-coil, Fig. 1, which is pointed towards the T3SS needle) is essential for its interaction with the needle protein PrgI^{16,30} led us to hypothesize that the distal region of SipD (defined here as the upper portion of the coiled-coil and mixed α/β domain, Fig. 1), is likely the site of interaction for the translocon. Here, we show results of NMR titrations between the tip proteins and the translocon proteins of Salmonella and Shigella. Our results show that the tip protein of both Salmonella and Shigella interacts with the N-terminal domain of their respective major translocon protein. The regions of the tip proteins involved in the interaction, the distal region, are conserved between the two bacteria. Additionally, SipD mutations at the distal region resulted in decreased invasion of Salmonella into human epithelial cells. Our results suggest a model of protein-protein interaction between the tip and the major translocon protein of the T3SS.

MATERIALS AND METHODS

Protein Expression and Purification

The expression and purification of SipD^{C244S} (residues 39-343 C244S) has been described previously.¹³ IpaD (residues 38-332) and IpaD^{C322S} (residues 38-332 C322S) were subcloned into the NdeI/SalI sites of a modified pET-21a expression vector containing an Nterminal His₆ tag followed by a tobacco etch virus (TEV) protease cleavage site. Both tip proteins retained a GHM artifact after cleavage with TEV protease. We previously reported that the C244S point mutation does not alter the crystal structure of SipD and Salmonella harboring the SipD^{C244S} mutation are fully functional in assembling the T3SS and invading eukaryotic cells.¹³ This suggests that Cys to Ser point mutations do not drastically alter the structure and function of T3SS tip proteins. SipB¹¹⁻²³², SipB⁸²⁻³¹², IpaB⁹⁻²²⁶ and IpaB⁷⁴⁻²²⁴ were subcloned into the NdeI/XhoI sites of pET-22b. The SipB and IpaB constructs retained a non-cleavable C-terminal LEH₆ tag for protein purification. Unlabeled SipB and IpaB proteins were expressed in E. coli BL21(DE3) DNAY cells in 1 liter of Luria Broth media containing 30 µg/mL kanamycin and 100 µg/mL carbenicillin. Cells were grown at 37°C until an OD₆₀₀ of ~0.8, induced with isopropyl- β -D-thiogalactopyrandoside (IPTG) (1 mM IPTG for SipB and 0.5 mM IPTG for IpaB) and grown overnight at 15°C to a final OD₆₀₀ of ~2.6. ILV labeling where the isoleucine $C\delta l$, the leucine $C\delta$, and the value $C\gamma$ methyl groups are ¹³C-labeled was used. Uniformly ¹⁵N/ILV-labeled tip proteins were expressed in 1 liter of 1× M9 minimal media at 37°C containing 1 g/L 15 NH₄Cl. At an OD₆₀₀ of ~ 0.4 the growth medium was supplemented with 60 mg of 2-ketobutyric acid-4-¹³C (Sigma #571342; which labels the isoleucine ${}^{13}C\delta$ 1 methyl group) and 100 mg of 2-keto-3-(methyl- ${}^{13}C$)butyric-4-¹³C acid (Sigma 571334; which labels the leucine ${}^{13}C\delta$ and the value ${}^{13}C\gamma$ methyl groups). For assignment of ILV 13C methyl resonances, perdeuterated 15N/ILVlabeled SipD^{C244S} was prepared by cell growth in M9 minimal media in 1 liter of D₂O with ¹⁵N- and ILV-labels, and 2 g/L deuterated D-glucose-1,2,3,4,5,6,6-d₇ (Cambridge Isotope Laboratories #CLM-2062). Protein expression was induced with 1 mM IPTG at an OD_{600} of 0.8 and cell growth was continued overnight at 15°C. Bacterial cells were harvested by centrifugation (4000 rpm, 10 min), resuspended with binding buffer (500 mM NaCl, 20 mM Tris-HCl, 5 mM imidazole, pH 8.0, 0.1 mM phenylmethanesulfonyl fluoride) and sonicated. The cell lysate was centrifuged (13,000 rpm, 10 min) to remove cellular debris and 700 µL of 5% polyethyleneimine was added to the supernatant to precipitate the nucleic acids and the mixture was centrifuged (13,000 rpm, 10 min). The supernatant was loaded into a 10 mL Ni²⁺-affinity column (Gold Biotechnology), and the column was washed with 150 mL binding buffer (500 mM NaCl, 20 mM Tris-HCl, 5 mM imidazole, pH 8.0), and eluted with 40 mL elution buffer (500 mM NaCl, 20 mM Tris-HCl, 250 mM imidazole, pH 8.0). For SipD and IpaD constructs, the eluted fractions were pooled and incubated in 0.02% by volume of 0.1 mM recombinant tobacco etch virus (TEV) protease³¹ in buffer (50 mM Tris-HCl, pH 8.0, 0.5 mM EDTA, 1 mM DTT, 20 mM NaCl) at room temperature overnight. The mixture was purified by Ni²⁺-affinity chromatography to separate the protein (which flowed through the Ni²⁺ column) from the affinity tag (which was retained in the Ni²⁺ column). Purified proteins were dialyzed into NMR buffer (100 mM NaCl, 20 mM sodium phosphate, pH 7.4 and 10% D₂O) and concentrated using Amicon Ultra 3K (Millipore) filtration columns. Protein concentrations were measured by A₂₈₀.

NMR Spectroscopy

NMR data were collected using a Bruker Avance 800 MHz spectrometer equipped with a cryogenic triple resonance probe and were processed using NMRPipe³² and analyzed using NMRView.³³ Two-dimensional ¹H-¹⁵N TROSY spectra were acquired at 30°C using ~0.4 mM of ¹⁵N/ILV SipD^{C244S} dissolved in NMR buffer with and without unlabeled SipB¹¹⁻²³² or SipB⁸²⁻³¹². Two-dimensional ¹H-¹⁵N-TROSY spectra were acquired at 30°C using ~0.2 mM ¹⁵N/ILV IpaD or ¹⁵N/ILV IpaD^{C322S} dissolved in NMR buffer with and without unlabeled IpaB⁹⁻²²⁶ or IpaB⁷⁴⁻²²⁴. Two IpaD constructs were used in the NMR titrations reported herein. Native IpaD has a cysteine residue at position 322, thus the NMR buffer also contained 5 mM dithiothreitol (DTT), whereas samples with IpaDC322S did not require DTT in the NMR buffer. The similarity of the 2D ¹H-¹⁵N TROSY spectra of IpaD and IpaD^{C322S} (Fig. S1) indicated that the C322S mutation did not drastically alter the tertiary structure of IpaD. IpaD^{C322S} was a better-behaved sample in solution with sharper NMR peaks and the sample was more stable over time, hence, IpaD^{C322S} was used in some of the titrations reported herein. The previously reported amide resonance assignments of SipD³⁴ and IpaD³⁵ were used in the NMR analysis. Residues perturbed during the titrations were mapped onto the crystal structure of SipD (PDB ID 3NZZ)¹³ or IpaD (PDB ID 2J00).¹⁵

Two-dimensional ¹H-¹³C HSQC spectra were acquired for SipD^{C244S} ILV-labeled samples with and without SipB. The ILV ¹³C-methyl resonances of SipD^{C244S} were assigned following the method of Xiao *et al.*³⁶ Briefly, 500 µL of ~0.9 mM perdeuterated ¹⁵N/ILV SipD^{C244S} in NMR buffer was lyophilized and resuspended in 100% D₂O. A 3D ¹H-¹³C-¹³C HMQC-NOESY-HMQC dataset was acquired using 8 scans with 2048 complex points (¹H), 80 complex points (¹³C) and 100 complex points (NOE ¹³C) with a 300 ms mixing time and a recycle delay of 2 s. Sweep widths were 10.0 ppm for ¹H centered at 4.69 ppm and 20.0 ppm for ¹³C centered at 19.0 ppm. To aid in the ILV assignment, twelve isoleucine to leucine point mutations were introduced into SipD^{C244S} using Quikchange (Stratagene). The Ile-to-Leu SipD^{C244S} mutants were expressed in M9 media and labeled at the isoleucine ¹³C δ I methyl group and 2D ¹H-¹³C HSQC spectra were acquired to identify the ¹³C δ I methyl peak. The rest of the leucine and valine ¹³C-methyl peaks were assigned from nuclear Overhauser effects (NOEs) from the 3D dataset in combination with distance information from the crystal structure of SipD (PDB ID 3NZZ).¹³

Salmonella Invasion Assay

A *Salmonella* invasion assay was used to test the effect of mutations at the distal end of SipD on the ability of *Salmonella typhimurium* to invade a cultured human intestinal epithelial cell line (Henle-407 cells, American Type Culture Collection CCL-6) as described previously.¹³ Wild type (SL1344) and *sipD*⁻ knockout strains of *Salmonella typhimurium* were used. Single, double, or triple point mutations were introduced by Quikchange kit (Stratagene) in the plasmid pRK2-SipD,¹³ which harbored full length *sipD*. Percent invasiveness was calculated relative to the *sipD*⁻ knockout strain complemented with the wild type SipD in pRK2-SipD. Error bars were determined from triplicate experiments.

Immunoblotting

Salmonella typhimurium strains with the sipD- knockout and pRK2-SipD (with wild type and mutant SipD) were grown in 15 mL of LB broth containing 0.3 M NaCl and 1 mM IPTG. Cell growth was continued at 37° C at 80 rpm until the OD₆₀₀ reached 0.9. Bacterial cells were harvested by centrifugation at 4000 rpm for 20 minutes then passed through a 0.45 µm pore-size filter. Proteins from the cultured supernatants were precipitated by incubating with 10% (v/v) trichloroacetic acid at 4° C for 1 hour and recovered by centrifugation at 4000 pm for 40 minutes. Pellets were resuspended in 2 mL 10% (v/v) SDS and 8 mL ice-cold acetone then incubated overnight at -20° C. After centrifugation at 4000 pm for 40 minutes, the pellets were dried and resuspended in 100 µL 8M Urea and 100 µL $2 \times$ SDS-loading dve. Proteins from whole-cell lysates (cell pellets) were resuspended in 1 mL 1× PBS and recovered by centrifugation at 13000 rpm for 15 minutes. The pellets were resuspended in 50 μ L 1× PBS and 50 μ L 2× SDS-loading dye. For immunoblotting, proteins were separated by SDS-PAGE and transferred to a PVDF membrane for 1.5 hours at 70 mA using an ECL Semi-Dry Blotter (Amersham Biosciences). After transfer, the membrane was incubated in 5% (v/v) non-fat dry milk with shaking for 1 hr at room temperature and then incubated with the primary antibody solution (1% non-fat dry milk, 0.1% Tween-20, $1\times$ TBS, rat anti-SipD IgG antibody) with shaking at 4°C overnight. Following this incubation step, the membrane was rinsed three times with wash buffer (0.1% Tween-20, $1 \times TBS$ pH 7.4). Membrane was then incubated in the secondary antibody solution (anti-rat IgG antibody conjugated to Alexa Fluor-680 in wash buffer) for 1 hr at room temperature with shaking. The membrane was then rinsed three times with wash buffer. The blot was analyzed using an ODYSSEY Infrared Imaging System (LI-COR Biosciences).

RESULTS

The N-terminal domain of SipB interacts with the distal region of SipD

We used NMR methods to identify the surfaces involved in the tip-translocon interaction by titrating the Salmonella SipB¹¹⁻²³² translocon protein into ¹⁵N/ILV labeled SipD^{C244S} and monitoring the titrations by acquiring 2D ¹H-¹⁵N TROSY spectra. Results of NMR titrations showed a concentration dependent decrease in intensities of many SipD^{C244S} peaks (Fig. 1A) indicating complex formation to be primarily in the intermediate exchange NMR time scale. In addition, some SipD^{C244S} residues, such as N211 (Fig. 1A), G220 and K302, and showed changes in the chemical shift with increasing concentrations of $SipB^{11-232}$ indicating these residues were in fast exchange NMR time scale. Importantly, there were also many peaks, such as A99, G176, S270, that did not undergo signal broadening, even at a SipD^{C244S}:SipB¹¹⁻²³² ratio of 1:2, indicating that complex formation did not result in protein aggregation, which would have broadened out all the peaks of SipD^{C244S}. To identify the SipD^{C244S} residues that were perturbed the most by SipB¹¹⁻²³², the peak intensity ratio $(I_{1,1}/I_{1,0})$ was calculated for each non-overlapped Sip^{DC244S} peak at a SipD^{C244S}:SipB¹¹⁻²³² molar ratio of 1:1 (Fig. 1B). Residues with peak intensity ratios lower than the average intensity minus one standard deviation were mapped onto the structure of SipD (Fig. 1C). SipD^{C244S} residues that were affected the most by SipB¹¹⁻²³² were primarily hydrophobic (L78, L116, A132, L150, G151, V156, L178, L240, L248, and L326) and some polar (S114, N211, N316) residues. The affected residues clustered in the distal

region of SipD (Fig. 1C), suggesting that this region of SipD is involved in the interaction with SipB.

The extreme N-terminal 11-27 residues of IpaB were previously reported to be essential for the interaction with IpaD.²³ Residues 11-27 of IpaB are homologous to residues 13-34 of SipB, and this region is present in the $SipB^{11-232}$ construct used in the NMR titration with SipD shown above (Fig. 1). To test if residues 11-34 of SipB are essential in the interaction with SipD, we used another construct, SipB⁸²⁻³¹², that lacked the N-terminal 13-34 residues for titrations with SipD^{C244S} and additionally contained residues 233–312, which are predicted to contain an additional a-helix. Titration of ¹⁵N/ILV SipD^{C244S} with unlabeled SipB⁸²⁻³¹² resulted in decreased peak intensities of specific SipD^{C244S} peaks (Fig. S2A) indicating that SipB⁸²⁻³¹² interacted with SipD^{C244S} despite lacking the N-terminal 13–34 residues. Plot of the peak intensity ratios of $(I_{1:1}/I_{1:0})$ of the SipD^{C244S}:SipB⁸²⁻³¹² complex at 1:1 molar ratio (Fig. S2B) and mapping the results on the structure of SipD showed that the affected residues of SipB⁸²⁻³¹² were located at the distal end of SipD (Fig. S2C). Again, many of the residues that were affected were hydrophobic or polar residues, such as L150, L194, V225, Y252 and Q297. There were some differences in the residues affected between the two different SipB constructs, however some residues were affected in both experiments, such as L150, D190 and N211 (Fig. 1B & Fig. S2B). These data suggested that SipD^{C244S} interacted with the N-terminal domain of SipB and that removal of the extreme N-terminal residues (11 to 81) of SipB did not significantly alter the affected SipD surface. Notably, we additionally attempted a titration between ¹⁵N/ILV SipD^{C244S} and SipB¹¹⁻³¹², but the NMR data was not acquired because sample precipitation was observed even at a SipD^{C244S}: SipB¹¹⁻³¹² molar ratio of 1:1.

Assignment of ILV ¹³C methyl resonances of SipD

Side-chain specific isotope-labeling techniques, such as the ¹³C methyl groups of isoleucine, leucine and valine (or ILV labeling), are utilized for NMR spectroscopy of high molecular weight proteins. Although ILV probes are useful in studying protein structures and dynamics because of their sharp NMR resonances and high sensitivity,³⁷ they have been sparsely used in studies of the T3SS tip proteins because of the current lack of ILV assignments for this protein family. The ¹H-¹³C HSQC spectra of ¹⁵N/ILV SipD^{C244S} showed 114 of 114 predicted methyl ILV resonances (Fig. S3). We report here that all the ILV ¹³C methyl resonances of SipD^{C244S} were assigned (Fig. S3) through a combination of site-directed mutagenesis of isoleucine to leucine (Fig. S4) and through-space ¹H-¹H nuclear Overhauser effects (NOEs) observed using 3D HMQC-NOESY of perdeuterated ILV SipD^{C244S} and distance information from the previously reported crystal structure of SipD (Fig. S5).¹³ These are currently the only available ILV assignments for any T3SS tip protein and should facilitate further studies of molecular interactions of SipD.

Titrations of ILV labeled SipD with SipB¹¹⁻²³²

The simultaneous labeling of SipD^{C244S} with ¹⁵N and ILV allowed the titration with SipB¹¹⁻²³² to be monitored firstly by 2D ¹H-¹⁵N HSQC to query the effect of the interaction on the ¹⁵N amides of SipD^{C244S}, the results of which are shown above (Fig. 1), and secondly by 2D ¹H-¹³C HSQC to query the effect of SipB¹¹⁻²³² on the ILV methyl resonances of

SipD^{C244S}. Like the results of the amide titrations (Fig. 1), most of the affected ILV resonances showed decreased peak intensities with a few residues showed chemical shift deviations, such as I46 (Fig. 2A) and I129. To identify which SipD^{C244S} ILV peaks were perturbed upon complex formation with SipB¹¹⁻²³², the (I_{1:1}/I_{1:0}) peak intensity ratio was plotted as a difference from peak intensity ratio of 1, which signified no change in the peak intensity upon complex formation (Fig. 2B). In agreement with the results of the amide titrations (Fig. 1C), the affected ILV methyl residues clustered at the distal end of SipD (Fig. 2C). Importantly, there were residues that were affected in ILV methyl titrations that were also affected in the amide titrations, such as V156, L178, V225, L240 (Fig. 1B & Fig. 2B). Unlike the ¹⁵N titrations, we observed the appearance of new slow-exchange methyl ILV peaks in the SipD^{C244S 1}H-¹³C HSQC spectra upon titration with SipB¹¹⁻²³² (Fig. 2A, arrows). The new peaks could be due to a conformational change in the side-chain methyls that was not observed in the ¹⁵N titrations. Together, the results of ILV methyl and amide titrations are in agreement that the distal region of SipD is the primary surface involved in the interaction with the N-terminal ectodomain of SipB.

The N-terminal domain of IpaB interacts with the distal end of IpaD

To compare the above results obtained for *Salmonella* on the interaction between the tip protein and the major translocon protein of *Shigella*, ¹⁵N/ILV labeled IpaD was titrated with unlabeled N-terminal domain of IpaB (residues 9-226). Results of NMR titrations monitored by 2D ¹H-¹⁵N TROSY showed that, similar to the *Salmonella* SipD and SipB interaction, stepwise addition of unlabeled IpaB⁹⁻²²⁶ to IpaD^{C322S} resulted in an IpaB⁹⁻²²⁶ concentration dependent decrease in peak intensities of specific IpaD^{C322S} amide peaks (Fig. 3A). Importantly, there were residues that did not undergo signal broadening, even at an IpaD^{C322S}:IpaB⁹⁻²²⁶ ratio of 1:2 indicating complex formation did not result in protein aggregation. The intensity ratio (I_{1:1}/I_{1:0}) for each non-overlapped IpaD^{C322S} residue at an IpaD^{C322S}:IpaB⁹⁻²²⁶ ratio of 1:1 was plotted to identify the tip residues that were perturbed the most upon addition of IpaB⁹⁻²²⁶ (Fig. 3B) and mapped onto the crystal structure of IpaD (Fig. 3C). Similar to the results observed for the *Salmonella* tip-translocon interaction, the affected residues clustered in the distal region of IpaD. IpaD^{C322S} residues that were perturbed the most by IpaB⁹⁻²²⁶ were primarily hydrophobic, such as Y160, V298 and I307, or polar residues, such as N58, E120 and K205 (Fig. 3B).

To test if the N-terminal region of IpaB (residues 11 to 27) was essential for the interaction between IpaD and IpaB as reported previously,²³ we titrated a shorter IpaB construct lacking the N-terminal region, IpaB⁷⁴⁻²²⁴, into IpaD. Titration of ¹⁵N/ILV IpaD with unlabeled IpaB⁷⁴⁻²²⁴ resulted in a concentration dependent decrease in peak intensity of many IpaD NMR peaks (Fig. S6A), indicating that IpaB interacted with IpaD despite lacking the Nterminal residues 11–27. Analysis of the peak intensity ratio (I_{1:1}/I_{1:0}) of IpaD:IpaB⁷⁴⁻²²⁴ complex at 1:1 molar ratio (Fig. S6B) showed that complex formation with IpaB⁷⁴⁻²²⁴ affected a cluster of IpaD residues at the distal region (Fig. S6C). Many of the IpaD residues that were affected were hydrophobic or charged polar residues, including R55, H131, I145, L195, Q239, M250 and L297 (Fig. S6B). There were some differences in the residues affected between the two different IpaB constructs, however many residues were affected in both experiments, such as E120, K205, S238, L260 and V298 (Fig. 3B & Fig. S6B).

Together, these data suggest that, like the *Salmonella* homologs, IpaD interacts with the N-terminal domain of IpaB and that removal of the extreme N-terminal residues of IpaB did not alter its interaction with IpaD.

The bile salt deoxycholate was reported to be essential for the interaction between IpaD and IpaB.²³ To test this, we titrated ¹⁵N labeled IpaD^{C322S} with unlabeled and IpaB⁹⁻²²⁶ in the presence of 0.4 mM deoxycholate. Like the titration in the absence of deoxycholate (Fig. 3), we observed a concentration dependent decrease in the peak intensity of IpaD^{C322S} NMR peaks (Fig. S7A). Analysis of the peak intensity ratio of IpaD^{C322S}:IpaB⁹⁻²²⁶ complex in the presence of deoxycholate complex at a 1:1:2 molar ratio (Fig. S7B) showed that the affected IpaD^{C322S} residues were primarily clustered at the distal region (Fig. S7C), similar to the results of the IpaD^{C322S}:IpaB⁹⁻²²⁶ complex in the absence of deoxycholate (Fig. 3C). IpaD residues that were affected in both the absence and presence of deoxycholate included Q148, Y160, Q220, W226 and I307 (Fig. 3B & Fig. S7B). Together, this suggests that deoxycholate did not largely alter the affected IpaD and IpaB interaction surface or modulate the protein affinity as observed by NMR.

Mutations in the distal end of SipD reduce invasion of *Salmonella* into human epithelial cells

To correlate the biological relevance of our NMR titration results, we tested the effect of mutations along the distal region of SipD on the ability of Salmonella to invade human intestinal epithelial Henle-407 cells using a previously reported *Salmonella* invasion assay.¹³ Deletion of the *sipD* gene rendered *Salmonella* completely non-invasive (Fig. 4A). However, invasiveness could be rescued by complementing the *sipD*⁻ null *Salmonella* strain with a plasmid expressing exogenous wild-type sipD (Fig. 4B). To test the significance of the distal end of SipD in invasion, point mutations were introduced into the mixed α/β region of the SipD rescue plasmid (Fig. 4B) and incorporated into the *sipD*⁻ strain. Point mutations either moderately reduced invasion (N196D and V191D) or had no effect (L271D and L280D) (Fig. 4A). Because protein-protein interactions often occur across large surfaces,³⁸ double or triple mutations in SipD were constructed (Fig. 4B). The double and triple mutations dramatically reduced the invasiveness of Salmonella (Fig. 4A). All mutant recombinant SipD showed similar protein fold to wild-type SipD based on circular dichroism (data not shown). Immunoblotting using anti-SipD antibodies of whole-cell lysates and supernatant showed that all the twelve SipD contructs used in the invasion assay were expressed in Salmonella, except for one triple mutant, N196D+V265D+L271D (Fig. S8). Thus, the decreased invasiveness of the SipD mutants (Fig. 4) could be ascribed to loss of function, except for one triple mutant, N196D+V265D+L271D, which was due to lost of protein expression (Fig. S8). These results suggest that the distal region of SipD is required for invasion and mutations that alter the surface of this region render Salmonella non-invasive.

DISCUSSION

The interaction of the tip protein and the major translocon protein is essential for the assembly of the T3SS needle apparatus. Dickenson *et al.*²³ showed by FRET and fluorescence polarization that the *Shigella* IpaD tip protein interacts with the N-terminal

ectodomain of the IpaB major translocon protein. Here, we used NMR methods to identify that the surfaces of the tip protein at the distal region, which includes the upper portion of the coiled-coil and the mixed α/β domain (Fig. 5C), are the sites of protein-protein interaction with the major translocon protein. The surfaces involved in the interaction are similar between SipD and SipB in Salmonella (Fig. 1 & Fig. 2) and IpaD and IpaB in Shigella (Fig. 3). Results of both amide (Fig. 1 & Fig. 3) and ILV titrations (Fig. 2) showed similar surfaces are involved in the interactions. Furthermore, in Salmonella the SipD residues affected were primarily hydrophobic and some uncharged polar residues (Fig. 1C), while in *Shigella* the IpaD residues affected were primarily charged polar and some hydrophobic residues (Fig. 3C). Changes in the NMR peaks brought about by proteinprotein interaction were primarily in the intermediate exchange time scale, suggesting dissociation constant near the μ M range (Fig. 1 & Fig. 3), which is in agreement with the reported K_d in the low µM range obtained by fluorescence spectroscopy.²³ Our results are in agreement with the proposed interaction surface for the tip and the major translocon protein identified by paramagnetic relaxation enhancement.³⁹ Additional data presented here show the role of deoxycholate and of the extreme N-terminal residues of the major translocon proteins in the interaction with the tip proteins; provides ILV methyl assignments of SipD; and provides biological invasion assay to test the proposed model of the tip-translocon interaction.

In contrast to the previous report that the bile salt deoxycholate is required for the IpaD-IpaB¹⁻²²⁶ interaction,²³ we observed IpaD-IpaB interaction in the absence of deoxycholate for the NMR titrations of IpaD-IpaB⁹⁻²²⁶ (Fig. 3) and IpaD-IpaB⁷⁴⁻²²⁴ (Fig. S6). Bile salts such as deoxycholate are secreted in the intestinal tract as part of the digestive system and they affect the type III secretion of the enteric bacteria *Shigella*^{40,41} and *Salmonella*.^{42,43} Deoxycholate binds to IpaD^{35,41} and SipD^{41,44}, and is hypothesized to function as an environmental sensor for the T3SS by inducing conformational change in IpaD that leads to latter steps in the assembly of the translocon.³⁵ It has been proposed that the interaction of deoxycholate with IpaD induces a conformational change at the distal region of IpaD which then promotes binding to IpaB.²³ We suspect that the difference in the results of fluorescence and NMR methods in observing IpaD-IpaB interaction with respect to the absence of deoxycholate lie in the concentrations of proteins used in the titrations. NMR requires 2-3 orders of magnitude more concentrated protein samples compared to fluorescence spectroscopy. Fluorescence polarization used 80 nM of fluorescein-labeled IpaB titrated with up to 10 µM of IpaD,23 whereas our NMR experiments used 0.2 mM IpaD titrated with up to 0.4 mM IpaB. Notably, we observed that the IpaD surface affected upon titration with IpaB was similar in the absence (Fig. 3C) and in the presence of 0.4 mM deoxycholate, or a 1:1:2 ratio of IpaD to IpaB to deoxycholate (Fig. S7C). Increasing the concentration of deoxycholate up to 1 mM, the concentration used by Dickenson et al.,23 resulted in sample precipitation.

Our results combined with previous data showing that the bottom of the coiled-coil of the tip protein is the site of protein-protein interactions with the needle protein¹⁶ leads to a model of the needle-tip-translocon interface. We propose a model of the needle-tip-translocon interface (Fig. 5) where the distal region of SipD interacts with the N-terminal ectodomain of SipB and the proximal end of SipD interacts with the needle PrgI. The *in vivo*

stoichiometry of the tip and translocon complex is poorly understood. However, one current hypothesis is that the tip complex either forms a homopentameric complex with five IpaD or heteropentameric complex consisting of one copy of IpaB and four copies of IpaD.⁹ Similarly, the *Salmonella* system is hypothesized to be composed to five SipD copies interacting with five PrgI needle subunits.¹⁶ Our NMR results that the major translocon protein interacts with the distal region of the tip protein agree with the recent model of the *Shigella* tip-translocon complex derived by electron microscopy and cross-linking of IpaD and IpaB.⁹ Our results are also in agreement with a recently published topology of the assembled major translocon protein AopB (a homolog of SipB and IpaB) from *Aeromonas hydrophila*.²⁹ Nguyen *et al*.²⁹ show that both the N and C-termini of AopB are extracellularly exposed when the translocon is assembled and therefore the N-terminal ectodomain of the major translocon protein is available for interaction with the tip protein.²⁹

CONCLUSIONS

We report here that the major translocon protein interacts with the distal surface of the tip protein in both *Salmonella* and *Shigella*. Mutations along the distal surface of the tip protein reduced the ability of *Salmonella* to invade intestinal epithelial cells, confirming the biological relevance of the distal surface of the tip protein. We also report the ILV methyl assignments of SipD and utilized the assignments in our interaction study to complement amide experiments (Fig. 2). Further structural studies are needed to determine which surface of the major translocon protein is affected upon interaction with the tip protein, whether the minor translocon protein is involved in the interaction with the tip and to build an atomic detail model of the assembled tip-translocon interface.

Data Deposition

NMR assignment of SipD^{C244S} ILV methyl resonances have been deposited into the BioMagResBank with the accession number 26739.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Figure 1.

Amide titrations of SipD^{C244S} with SipB¹¹⁻²³². (A) Overlay of four ¹H-¹⁵N TROSY spectra of ¹⁵N/ILV SipD^{C244S} with increasing concentrations of unlabeled SipB¹¹⁻²³². (B) Plot of ratio of peak intensities of complex (I_{1:1}) vs free (I_{1:0}) for the SipD^{C244S}:SipB¹¹⁻²³² complex at 1.1:1.0 molar ratio shown with the average peak intensity ratio (dashed gray line) and one standard deviation from the average (1 σ , dashed red line). (C) Residues (colored red) with peak intensity ratio (I_{1:1}/I_{1:0}) lower than 1 σ were mapped onto the crystal structure of SipD [colored green (N-terminal hairpin), gray (coiled-coil), and orange (mixed α/β domain)].



Figure 2.

ILV titrations of SipD^{C244S} with SipB¹¹⁻²³². (A) Overlay of four ¹H-¹³C HSQC spectra of ¹⁵N/ILV SipD^{C244S} with increasing concentrations of unlabeled SipB¹¹⁻²³². (B) Plot of peak intensities subtracted from one $(1 - I_{1:1}/I_{1:0})$ of the SipD^{C244S}:SipB¹¹⁻²³² complex at molar ratio of 1.1:1.0 shown with the average value (dashed gray line) and one standard deviation from the average (1 σ , dashed red line). (C) Residues (colored red) with (1 - I_{1:1}/I_{1:0}) value higher than 1 σ were mapped onto the crystal structure of SipD [colored green (N-terminal hairpin), gray (coiled-coil), and orange (mixed α/β domain)].



Figure 3.

Amide titrations of IpaD^{C322S} with IpaB⁹⁻²²⁶. (**A**) Overlay of four ¹H-¹⁵N TROSY spectra of ¹⁵N/ILV IpaD^{C322S} with increasing concentrations of unlabeled IpaB⁹⁻²²⁶. (**B**) Plot of ratio of peak intensities of complex (I_{1:1}) vs free (I_{1:0}) for the IpaDC^{322S}:IpaB⁹⁻²²⁶ complex at 1.1:1.0 molar ratio shown with the average peak intensity ratio (dashed gray line) and one standard deviation from the average (1 σ , dashed red line). (**C**) Residues (colored red) with peak intensity ratio (I_{1:1}/I_{1:0}) lower than 1 σ were mapped onto the crystal structure of IpaD [colored green (N-terminal hairpin), gray (coiled-coil), and orange (mixed α/β domain)].

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Figure 4.

Mutations in the distal region of SipD decrease invasion of *Salmonella*. (A) Deletion of *sipD* (*sipD*) renders *Salmonella* non-invasive to Henle-407 human intestinal epithelial cells. Invasion was rescued by complementing with a plasmid containing wild-type *sipD*. Point mutations in the mixed α/β domain of SipD reduced invasion (N196D and V191D) or had no effect (L271D and L280D). Double and triple mutations decreased invasion even further. (B) The residues chosen for mutation are shown in red on the structure of SipD.



Figure 5.

Proposed model of the needle, tip and translocon interface. (A) Membrane topology of SipB showing the N-terminal domain (NTD). (B) Structure of the SipB N-terminal domain (82–226) with predicted secondary structure of regions of unknown structure present in our SipB constructs. (C) Model for the interaction hubs on the tip protein SipD, where the N-terminal domain of the translocon SipB interacts with the mixed α/β domain and the needle PrgI in interacts with the bottom of the coiled-coiled. The model of the *Shigella* IpaD and IpaB is expected to be homologous.